Assignment 2 - Solving two 1D problems

This task is also hosted here - https://github.com/jhenderson0/jh_ucl_HPC/tree/main/Task%202 (https://github.com/jhenderson0/jh_ucl_HPC/tree/main/Task%202

```
In [37]: import numpy as np
%matplotlib inline
    from matplotlib import pyplot as plt
    from matplotlib import cm
    import seaborn as sns
    plt.style.use('seaborn-notebook')
```

Part 1: Solving a wave problem with sparse matrices

In this part of the assignment, we want to compute the solution to the following (time-harmonic) wave problem:

$$\frac{d^2 u}{dx^2} + k^2 u = 0 \qquad \text{in } (0,1),$$

$$u = 0 \qquad \text{if } x = 0,$$

$$u = 1 \qquad \text{if } x = 1,$$

with wavenumber $k = 29\pi/2$.

In this part, we will approximately solving this problem using the method of finite differences. We do this by taking an evenly spaced values $x_0 = 0, x_1, x_2, \dots, x_N = 1$ and approximating the value of u for each value: we will call these approximations u_i . To compute these approximations, we use the approximation

$$\frac{\mathrm{d}^2 u_i}{\mathrm{d}x^2} \approx \frac{u_{i-1} - 2u_i + u_{i+1}}{h^2},$$

where h = 1/N.

With a bit of algebra, we see that the wave problem can be written as

$$(2 - h^2 k^2) u_i - u_{i-1} - u_{i+1} = 0$$

if x_i is not 0 or 1, and

$$u_i = 0$$
 if $x_i = 0$,
 $u_i = 1$ if $x_i = 1$.

This information can be used to re-write the problem as the matrix-vector problem $A\mathbf{u} = \mathbf{f}$, where A is a known matrix, \mathbf{f} is a known vector, and \mathbf{u} is an unknown vector that we want to compute. The entries of \mathbf{f} and \mathbf{u} are given by

$$[\mathbf{u}]_i = u_i,$$

$$[\mathbf{f}]_i = \begin{cases} 1 & \text{if } i = N, \\ 0 & \text{otherwise.} \end{cases}$$

The rows of A are given by

$$[A]_{i,j} = \begin{cases} 1 & \text{if } i = j, \\ 0 & \text{otherwise,} \end{cases}$$

if i = 0 or i = N; and

$$[A]_{i,j} = \begin{cases} 2 - h^2 k^2 & \text{if } j = i, \\ -1 & \text{if } j = i+1, \\ -1 & \text{if } j = i-1. \\ 0 & \text{otherwise,} \end{cases}$$

otherwise.

a)

Write a Python function that takes N as an input and returns the matrix A and vector f. You should use an appropriate sparse storage format for the matrix A.

```
In [38]: from scipy.sparse import coo_matrix
```

```
In [39]: def discretise wave(N):
             """Generate the matrix and rhs associated with the discrete time-harmonic wave equatio
             h = 1/N
             k = 29*np.pi/2
             #Number of elements is N+1 for diagonal + the off diagonal (2N) - the two zeros on the
          top and bottom row
             nelements = N+1 + 2*N - 2
             #Set up coo styled data storage
             row_ind = np.zeros(nelements, dtype=int)
             col ind = np.zeros(nelements, dtype=int)
             data = np.zeros(nelements, dtype=np.float64)
             #Create the vector (all zeros with a 1 at the end)
             f = np.concatenate((np.zeros(N, dtype=np.float64),[1]))
             #Set the 0,0 element to be 1
             row_ind[0] = col_ind[0] = 0
             data[0] = 1
             #Set the N,N element to be 1
             row_ind[-1] = col_ind[-1] = N
             data[-1] = 1
             #Loop through each row
             count = 1
             for i in range(1,N):
               #For each row we have three data entries
               row_ind[count : count+3] = i
               #For the diagonals we have 2-h^2k^2
               col_ind[count] = i
               data[count] = 2-h**2*k**2
               #For the off diagonals we have -1
               col ind[count + 1] = i + 1
               col_ind[count + 2] = i - 1
               data[count + 1 : count + 3] = -1
               count += 3
             return coo_matrix((data, (row_ind, col_ind)), shape=(N+1, N+1)).tocsr(), f
```

A quick example, if we use N = 2 we expect to get a matrix out which looks like:

$$\mathbf{A} = \begin{bmatrix} 1 & 0 & 0 \\ -1 & 2 - h^2 k^2 & -1 \\ 0 & 0 & 1 \end{bmatrix}$$

And our vector should be:

$$\mathbf{f} = \begin{bmatrix} 0 \\ 0 \\ 1 \end{bmatrix}$$

b)

The function <code>scipy.sparse.linalg.spsolve</code> can be used to solve a sparse matrix-vector problem. Use this to **compute the** approximate solution for your problem for N=10, N=100, and N=1000. Use <code>matplotlib</code> (or any other plotting library) to plot the solutions for these three values of N.

```
In [41]: from scipy.sparse.linalg import spsolve
In [42]:
           sols = []
           Ns = [10, 100, 1000]
           for N in Ns:
             A, f = discretise_wave(N)
             sols.append(spsolve(A, f))
In [43]: fig, ax = plt.subplots(1,3,figsize=(15, 5),dpi=100)
           for idx, N in enumerate(Ns):
             x = np.linspace(0,1,N+1)
             ax[idx].plot(x,sols[idx],c='b')
             ax[idx].set_ylabel("u(x)")
             ax[idx].set_xlabel("x")
             ax[idx].set_title(f"N = {N}")
                                                               N = 100
                                                                                                   N = 1000
                             N = 10
             1.0
                                                                                    1.00
                                                 1.0
                                                                                   0.75
             0.8
                                                                                    0.50
                                                 0.5
             0.6
                                                                                   0.25
                                                0.0
                                                                                   0.00
             0.4
                                                                                   -0.25
                                                -0.5
             0.2
                                                                                   -0.50
                                                                                   -0.75
             0.0
                                                -1.0
                                                                                   -1.00
                 0.0
                      0.2
                                 0.6
                                      0.8
                                           1.0
                                                    0.0
                                                         0.2
                                                                    0.6
                                                                               1.0
                                                                                        0.0
                                                                                                                   1.0
```

c)

Briefly (1-2 sentences) comment on your plots: How different are they to each other? Which do you expect to be closest to the actual solution of the wave problem?

Answer - The N=10 solution is very different to the other two - it has only managed to really meet the boundary conditions and not produced a wave. The other two solutions are closer to eachother, both show the same overall wave. The greater the value of N used, the smoother the function becomes and the closer I would expect it to be to the true analytical solution.

d)

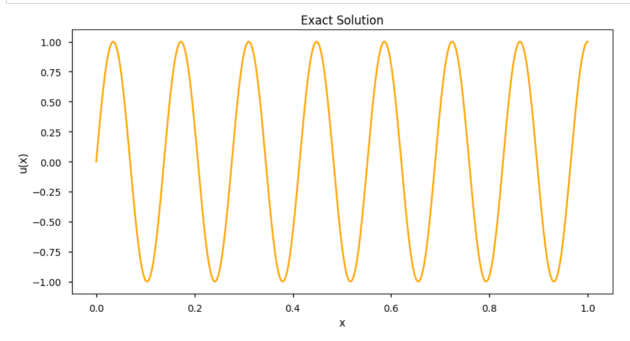
This wave problem was carefully chosen so that its exact solution is known: this solution is $u_{\text{exact}}(x) = \sin(kx)$. (You can check this by differentiating this twice and substituting, but you do not need to do this part of this assignment.)

A possible approximate measure of the error in your solution can be found by computing $\max_i |u_i - u_{\text{exact}}(x_i)| \; .$

Compute this error for a range of values for N of your choice, for the methods you wrote in both parts 1 and 2. On axes that both use log scales, plot N against the error in your solution. You should pick a range of values for N so that this plot will give you useful information about the methods.

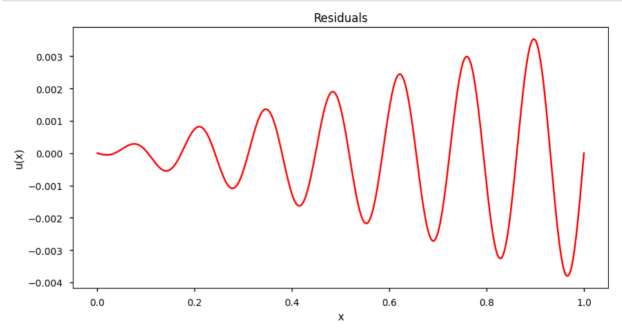
A plot to see the exact solution:

```
In [45]: fig, ax = plt.subplots(1,figsize=(10, 5),dpi=100)
    x = np.linspace(0,1,1001)
    ax.plot(x,exact(x),c='orange')
    ax.set_ylabel("u(x)")
    ax.set_xlabel("x")
    ax.set_title("Exact Solution");
```



A plot of the residuals between the solved values and exact values:

```
In [46]: fig, ax = plt.subplots(1,figsize=(10, 5),dpi=100)
    x = np.linspace(0,1,1001)
    ax.plot(x,exact(x)-sols[2],c='red')
    ax.set_ylabel("u(x)")
    ax.set_xlabel("x")
    ax.set_title("Residuals");
```



Next we define the error function:

```
In [47]: def err(u,u_exact):
    return np.amax(np.absolute(u-u_exact))
```

And then check the error for increasing N:

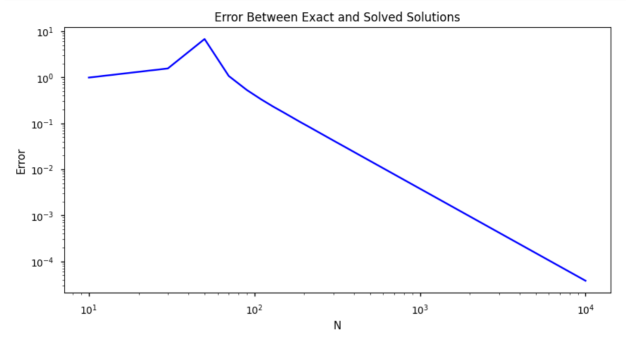
```
In [48]: Ns = np.linspace(10,10000,500,dtype=int)
errs = np.zeros(len(Ns))
for idx, N in enumerate(Ns):

#Solve the equations
A, f = discretise_wave(N)
sol = spsolve(A, f)

#Calcualte the exact value
ex = exact(np.linspace(0,1,N+1))

#Calcualte the error
errs[idx] = err(sol,ex)
```

```
In [49]: fig, ax = plt.subplots(1,figsize=(10, 5),dpi=100)
    ax.plot(Ns,errs,c='b')
    ax.set_ylabel("Error")
    ax.set_xlabel("N")
    ax.set_yscale("log")
    ax.set_yscale("log")
    ax.set_title("Error Between Exact and Solved Solutions");
```

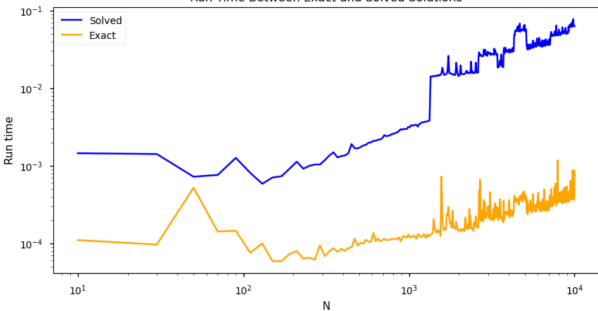


e)

For the same values of N, measure the time taken to compute your approximations for both functions. On axes that both use \log scales, plot N against the time taken to compute a solution.

```
In [54]: fig, ax = plt.subplots(1,figsize=(10, 5),dpi=100)
    ax.plot(Ns,times[0],label = "Solved",c='blue')
    ax.plot(Ns,times[1] , label = "Exact",c='orange')
    ax.set_ylabel("Run time")
    ax.set_xlabel("N")
    ax.set_yscale("log")
    ax.set_xscale("log")
    ax.set_title("Run Time Between Exact and Solved Solutions")
    ax.legend();
```





f)

We now want to compute an approximate solution where the measure of error is 10^{-8} or less. By looking at your plots, pick a value of N that you would expect to give error of 10^{-8} or less. Briefly (1-2 sentences) explain how you picked your value of N and predict how long the computation will take.

Answer - I would choose a value of N of the order of maginutude $\approx 10^6$. This is because the order of maginutude of the error roughly decreases linearly with the order of magnitude of N such that:

$$\log(\text{Err}(N)) \approx -\frac{1}{2}\log(N) + c$$

As for run time, it seems like this is linked to N roughly by:

$$\log(\operatorname{Time}(N)) \approx \log(N) + d$$

Where c and d are constants. I would expect a run time around order of maginutude 10^0 seconds

g)

Compute the approximate solution with your value of N. Measure the time taken and the error, and briefly (1-2 sentences) comment on how these compare to your predictions.

```
In [55]: def time and error(N):
           #Solve the equations
           start = time()
           A, f = discretise_wave(N)
           sol = spsolve(A, f)
           run_time = time() - start
           #Calculate the exact value
           ex = exact(np.linspace(0,1,N+1))
           #Calculate the error
           er = err(sol, ex)
           print(f"N: {N}")
           print(f"Error: {er}")
           print(f"Runtime: {run time}")
In [56]: time_and_error(int(1e6))
```

N: 1000000 Error: 1.5517885523438912e-08

Runtime: 2.520681619644165

Answer - The error isn't quite as low as I predicted - I expected this as my estimates were only made very crudely by checking how many orders of maginitude the runtime or the error changed when I changed the order of magnitude of N - however, the order of magintude for both the error and run time are as I expected. Interestingly, if N is increased very slightly from 10^6 , which I would expect to bring down the error, the error instead increases while the runtime continues with the trend of increasing:

```
In [57]: | time_and_error(int(1.00001e6))
         time_and_error(int(1e7))
         N: 1000010
         Error: 2.0754513803400432e-07
         Runtime: 2.4040367603302
         N: 10000000
         Error: 0.00011509646200105403
         Runtime: 25.023558855056763
```

Part 2: Solving the heat equation with GPU acceleration

In this part of the assignment, we want to solve the heat equation

$$\frac{du}{dt} = \frac{1}{1000} \frac{d^2 u}{dx^2} \quad \text{for } x \in (0, 1),$$

$$u(x, 0) = 0,$$

$$u(0, t) = 10,$$

$$u(1, t) = 10.$$

This represents a rod that starts at 0 temperature which is heated to a temperature of 10 at both ends.

Again, we will approximately solve this by taking an evenly spaced values $x_0 = 0, x_1, x_2, \dots, x_N = 1$. Additionally, we will take a set of evenly spaced times $t_0 = 0, t_1 = h, t_2 = 2h, t_3 = 3h, \dots$, where h = 1/N. We will write $u_i^{(j)}$ for the approximate value of u at point x_i and time t_i (ie $u_i^{(j)} \approx u(x_i, t_i)$).

Approximating both derivatives (similar to what we did in part 1), and doing some algebra, we can rewrite the heat equation as

$$u_i^{(j+1)} = u_i^{(j)} + \frac{u_{i-1}^{(j)} - 2u_i^{(j)} + u_{i+1}^{(j)}}{1000h},$$

$$u_i^{(0)} = 0,$$

$$u_0^{(j)} = 10,$$

$$u_N^{(j)} = 10.$$

This leads us to an iterative method for solving this problem: first, at t = 0, we set

$$u_i^{(0)} = \begin{cases} 10 & \text{if } i = 0 \text{ or } i = N, \\ 0 & \text{otherwise;} \end{cases}$$

then for all later values of time, we set

$$u_i^{(j+1)} = \begin{cases} 10 & \text{if } x = 0 \text{ or } x = N, \\ u_i^{(j)} + \frac{u_{i-1}^{(j)} - 2u_i^{(j)} + u_{i+1}^{(j)}}{1000h} & \text{otherwise.} \end{cases}$$

a)

Implement this iterative scheme in Python. You should implement this as a function that takes N as an input.

```
In [58]: import numba
```

My function below takes the temperature values of a rod, u(x) discretised into N+1 indexes, and updates them by iterating through the heat equation up to some maximum time, t_{max} :

```
In [59]: @numba.njit(['int32,int32,float64[:,:]'])
def solve_heat(N,tmax,u):

#Loop through times
for j in range((N*tmax)):
    #Update the points in the rod excluding the end points
    u[j+1,1:N] = u[j,1:N] + (u[j,0:N-1] - 2*u[j,1:N] + u[j,2:N+1])* N/1000
```

If memory was an issue (in the case of a large t_{max}) I would have the function above only update a 1D N+1 long array and not store the rod values for each timestep. As memory is not too much of an issue in this scenario, I save all of the time step data points as this makes plotting simple:

```
In [60]: N = 3
          tmax = 2
          #Create the rod and set the ends to a constant temperature
          u = np.zeros((N*tmax+1,N+1))
          u[:,0] = u[:,-1] = 10
          solve_heat(N,tmax,u)
Out[60]: array([[10.
                        , 0.
                                              0.
                                                         , 10.
                                                                      ],
                           , 0.03 , 0.03 , 10.
, 0.05991 , 0.05991 , 10.
                 [10.
                                                                      ],
                 [10.
                                                                      ],
                            , 0.08973027, 0.08973027, 10.
                 [10.
                                                                      ],
                            , 0.11946108, 0.11946108, 10.
                 [10.
                                                                      ],
                           , 0.1491027 , 0.1491027 , 10.
, 0.17865539, 0.17865539, 10.
                 [10.
                                                                      ],
                 [10.
                                                                      ]])
```

b)

Using a sensible value of N, plot the temperature of the rod at t=1, t=2 and t=10. Briefly (1-2 sentences) comment on how you picked a value for N.

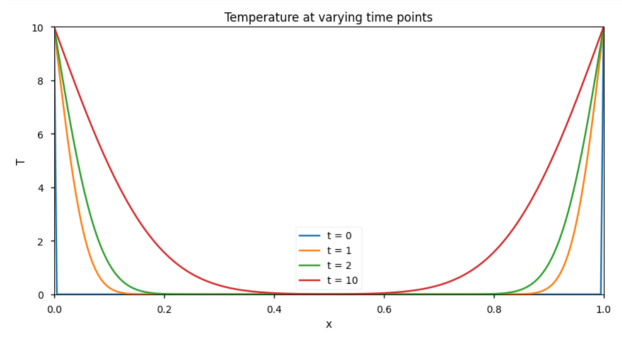
Answer - I have used a value of N which provides me with enough x values to give a smooth plot while being low enough to run quickly:

Below is a plot showing the temperature across the rod for different time values:

```
In [62]: fig, ax = plt.subplots(1,figsize=(10, 5),dpi=100)

ts = [0,1,2,10]
    for t in ts:
        ax.plot(np.linspace(0,1,N+1),u[int(t*N)],label=f't = {t}')

ax.set_ylabel("T")
        ax.set_xlabel("x")
        ax.set_xlim([0,1])
        ax.set_ylim([0,10])
        ax.set_title("Temperature at varying time points")
        ax.legend();
```



A heatmap visulisation of the rod:

```
In [63]: fig, ax = plt.subplots(4,1,figsize=(10, 5))
          for idx,t in enumerate(ts):
            \verb|sns.heatmap(u[int(t*N)][np.newaxis,:], cmap='hot',ax=ax[idx]|)|
            ax[idx].set_xlabel("x")
            ax[idx].set_ylabel(f"t = {t}")
            ax[idx].set_yticks([])
            ax[idx].set_xticks([])
          fig.tight_layout()
                                                                                           - 10
           t = 0
                                                                                           - 5
                                                                                           - 10
                                                                                           - 5
                                                                                           - 10
                                                                                           - 5
                                                                                           - 10
           t = 10
                                                                                           - 5
```

Use numba.cuda to parallelise your implementation on a GPU. You should think carefully about when data needs to be copied, and be careful not to copy data to/from the GPU when not needed.

We want to parallelise the process of updating each point on the bar:

```
In [65]: #Our kernel function - iterates through the bar
@cuda.jit
def solve_heat_cuda(u0,u1):

    # Index of thread on GPU
    i = cuda.grid(1)

    #Get N from the size of the array
    N = u0.size-1

#Update the points in the rod excluding the end points
if(i > 0 and i < N):
    #Update the vector for the next time point
    u1[i] = u0[i] + (u0[i + 1] - 2*u0[i] + u0[i - 1])*N/1000</pre>
```

```
In [68]: N = 3
         tmax = 2
         #Create the rod and set the ends to a constant temperature
         u = np.zeros((N+1))
         u[0] = u[-1] = 10
         # Manage the number of threads
         threadsperblock = 32
         blockspergrid = (u.size + (threadsperblock - 1)) // threadsperblock
         #Add the rod to the GPU plus a buffer to be updated
         u0 = cuda.to_device(u)
         u1 = cuda.to_device(u)
         #Update the rod
         for j in range(N*tmax):
           if (j % 2) == 0:
             solve heat cuda[blockspergrid, threadsperblock](u0,u1)
             solve heat cuda[blockspergrid, threadsperblock](u1,u0)
         #Copy the final timestep data back to us
         u = u0.copy_to_host()
         print(u)
         [10.
                       0.17865539 0.17865539 10.
                                                          ]
```

```
In [69]: | N = 300
        tmax = 200
        ##CPU BASED##
        uCPU = np.zeros((N*tmax+1,N+1))
        uCPU[:,0] = uCPU[:,-1] = 10
        start = time()
        solve_heat(N,tmax,uCPU)
        print(f"CPU Runtime: {time()-start}")
        ##GPU BASED##
        uGPU = np.zeros((N+1))
        uGPU[0] = uGPU[-1] = 10
        # Manage the number of threads
        threadsperblock = 32
        blockspergrid = (uGPU.size + (threadsperblock - 1)) // threadsperblock
        #Add the rod to the GPU
        u0 = cuda.to device(uGPU)
        u1 = cuda.to_device(uGPU)
        start = time()
        #Update the rod
        for j in range(N*tmax):
          if (j % 2) == 0:
            solve_heat_cuda[blockspergrid, threadsperblock](u0,u1)
            solve_heat_cuda[blockspergrid, threadsperblock](u1,u0)
        #Ensure the GPU code is finished before we get the end time
        cuda.synchronize()
        print(f"GPU Runtime: {time()-start}")
        print(f"Total number of threads: {threadsperblock*blockspergrid}")
        #Copy the final timestep data back to us
        uGPU = u0.copy_to_host()
        print(f"Function difference: {np.sum(uGPU-uCPU[-1])}")
        np.testing.assert_allclose(uGPU,uCPU[-1])
        CPU Runtime: 0.049661874771118164
        GPU Runtime: 3.397979497909546
```

d)

Total number of threads: 320 Function difference: 0.0

Use your code to estimate the time at which the temperature of the midpoint of the rod first exceeds a temperature of 9.8. Briefly (2-3 sentences) describe how you estimated this time. You may choose to use a plot or diagram to aid your description, but it is not essential to include a plot.

Answer - For this question I used my CPU based code which returns to me all values of the rod up to a maximum time t_{max} . I then check to see at what time step the mid point value first exceeded 9.8 and return that timestep converted into seconds:

```
In [70]: def find_time(u,T):
    N = u.shape[1] - 1

#Find all the mid points
    mid_points = u[:,int((N+1)/2)]

try:
    #Get the first mid point value which goes above T - convert from timestep into seconds
    timeT = np.asarray(mid_points > T).nonzero()[0][0]/N
    print(f"Mid point temperature > {T} found at t = {timeT}")
    return timeT

except:
    print(f"Mid point temperature > {T} not found")
```

```
In [71]: #Run the heat equation for a large max time values
    N = 400
    tmax = 450

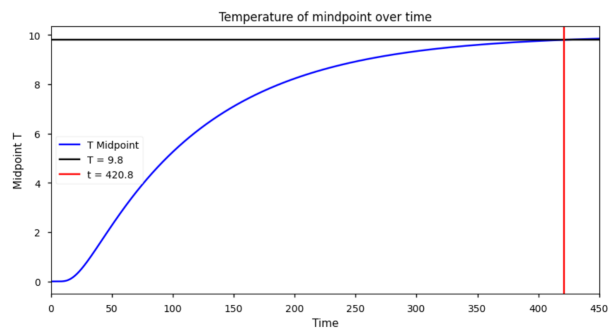
#Create the rod and set the ends to a constant temperature
    u = np.zeros((N*tmax+1,N+1))
    u[:,0] = 10
    u[:,-1] = 10

#Solve the heat equation
    solve_heat(N,tmax,u)

#Check to see if we've found a midpoint temperature of 9.8
    timeT = find_time(u,9.8)
```

Mid point temperature > 9.8 found at t = 420.845

```
In [72]: fig, ax = plt.subplots(1,figsize=(10, 5),dpi=100)
    ax.plot(np.linspace(0,tmax,N*tmax+1),u[:,int((N+1)/2)],label = 'T Midpoint',c='b')
    ax.set_ylabel("Midpoint T")
    ax.set_xlabel("Time")
    ax.set_title("Temperature of mindpoint over time")
    ax.set_xlim([0,tmax])
    ax.axhline(9.8,c='black',label = 'T = 9.8')
    ax.axvline(timeT,c='red',label = f't = {timeT:0.1f}')
    ax.legend();
```



Answer Continued - If I was to do this using my GPU based code, I would run the time stepping loop until a midpoint temperature of 9.8 was identified. I could also use my array of CPU generated mid point values and then create an array with an equal number of points going from 0 to t_{max} , interpolation could then be used to identify the time at which the temperature first exceeded 9.8.

In [73]: print(f"Interpolated time: {np.interp(9.8, u[:,int((N+1)/2)], np.linspace(0,tmax,N*tmax+1))}
}")

Interpolated time: 420.8428504064262